Nanocomposite Catalysts for Steam Reforming of Methane and Biofuels: Design and Performance

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1. Introduction

Design of nanocomposite materials with high mixed ionic-electronic conductivity (MIEC) and oxygen mobility possessing a high and stable performance in real operation conditions is now considered as one of the most promising trends in developing new anode materials for IT SOFC (Primdahl & Mogensen, 2002; Atkinson et al., 2004; Wincewicz & Cooper, 2005; Dicks, 1998; Kharton et al., 2006; Marina et al., 1999; Xia & Liu, 2002; Zha et al., 2004; Ishihara et al., 2000) and structured catalysts for steam/autothermal reforming of gas and liquid fuels (Souza & Schmal, 2003; Domine et al., 2008; Sadykov et al., 2009).

State-of-the art Ni/Y2O3–ZrO2 (Ni/YSZ) cermet anodes of solid oxide fuel cells (SOFC) have excellent catalytic properties and stability in the oxidation of hydrogen fuel at SOFC operation conditions (Atkinson et al., 2004). However, the lack of a hydrogen infrastructure and the unsolved hydrogen storage problem have initiated the research aimed at direct utilization of natural gas, which represents one of the key aspects of SOFC technology. Internal steam reforming (SR) is the most promising concept in using the natural gas (as well as bio-gas or bioethanol) as a fuel (Wincewicz & Cooper, 2005; Dicks, 1998). In this case, the reaction takes place directly in the anode compartment, allowing a better management within the stack of heat produced by the exothermic electrochemical oxidation and consumed by the endothermic reforming reaction. Unfortunately, with the Ni/YSZ cermet,
coking occurs leading to the deterioration of anode performance (Atkinson et al., 2004; Dicks, 1998). Under high carbon activity environment, Ni metal could also be corroded by the metal dusting. Ni/YSZ cermet anodes can only be used in hydrocarbon fuels if excess steam is present to suppress the carbon deposition, which, however, decreases the electrical efficiency of the cells (Winczewicz & Cooper, 2005; Dicks, 1998). Hence, development of robust anodes with a high and stable activity in the internal reforming of fuels is vital for the natural gas/biogas/bioethanol fuel-based SOFC. Next approaches were suggested up to date to solve this problem:

1. Partial or complete replacement of doped zirconia with doped ceria or ceria-zirconia possessing a higher lattice oxygen mobility/reactivity (Atkinson et al., 2004; Winczewicz & Cooper, 2005; Dicks, 1998; Kharton et al, 2006; Marina et al, 1999; Xia & Liu, 2002; Zha et al, 2004; Ishihara et al, 2000). This is combined with partial or complete replacement of Ni by copper (Dongare et al., 2002; Ana et al., 2004). This approach allows not only to prevent coking but also ensures a high performance in SR at intermediate temperatures. In this case, some problems could be caused by considerable chemical expansion of ceria lattice in strongly reducing conditions as well as by copper sintering.

2. Replacing doped zirconia by perovskites (mainly, doped La chromites) promoted by metals active in SR (Ni or precious metals, mainly, Ru) (Peña-Martínez et al., 2006; Sfeir et al., 2001; Sauvet & Irvine, 2004; Vernoux et al., 1998; Wan & Goodenough, 2005). In this case, insufficient conductivity of perovskites in reducing conditions could be a problem (Plint et al., 2006).

3. Partial or complete replacement of Ni in Ni/YSZ or Ni/perovskite cerments by precious metals (Wan & Goodenough, 2005; Bebelis et al., 2006; Suzuki et al., 1993; Takeguchi et al., 2003). This allows to suppress coking and enhance the middle-temperature performance which is explained by a much higher specific catalytic activity of precious metals in CH₄ SR, Pt being the most active metal (Wei & Iglesia, 2004). In the case of cerments containing only precious metals, apparent drawback is their high cost.

Hence, the most promising approach for achieving a high level of anode activity at middle temperatures in CH₄ steam reforming while preventing coking, keeping a high level of conductivity and a low cost, is to promote Ni/YSZ(ScSZ) cerments by fluorite-like (doped ceria-zirconia) or perovskite-like (mixed chromates-manganites) oxides along with small (~1%) amounts of precious metals (Pt, Pd, Ru). These (nano)composites are comprised of components able to efficiently activate C-H and C-C bonds in the fuel molecules (Ni, precious metals) and oxide components providing activation of water molecules and transfer of hydroxyls and/or hydroxocarbonate/oxygen species to the metal particles where they interact with activated C-H-O species producing syngas (Souza & Schmal, 2003).

In the last years biomass has been recognized as one of the major world renewable energy sources. Bio-oil derived from the fast pyrolysis of biomass or bio-ethanol can be converted via steam reforming into hydrogen or syngas, which can be further used in fuel cells or directed to synthesis of liquid fuels and valuable chemicals (Asadullah et al., 2002). For SOFC, an attractive option is direct internal reforming of bio-fuels on catalytically active anodes (Jamsak et al., 2007). Hence, efficient, inexpensive and robust catalysts for the steam reforming of biofuel are required. The most demanding problem of their design is a heavy coking of catalysts even in the feeds with the excess of steam caused by a high reactivity of bio-fuel components (carboxylic acids, aldehydes, ketones, alcohols etc), thus excluding application of traditional Ni-based steam reforming catalysts, Ni-YSZ anode cermets (Jamsak et al., 2007) or precious metals (Pt, Pd, Ru) supported on alumina, zirconia, etc.
Haryanto et al., 2005; Breen et al., 2002). Hence, inexpensive nanocomposite catalysts on the base of promoted Ni/could be also attractive as active and stable to coking components of catalysts for steam reforming of biofuels.

To ensure a high performance of these composites in steam reforming of a given type of fuel, their composition and preparation procedures are to be properly optimized. Specificity of functional characteristics of nanocomposite materials strongly depends on the properties of interfaces/domain boundaries which could act as paths for fast oxygen diffusion and generate specific surface sites responsible for activation of reagents. Chemical composition and local structure of these interfaces controlling their properties are determined both by the nature of coexisting phases and their interaction depending upon the size of domains, their disordering and nanocomposite synthesis procedure.

This chapter overviews results of research within the frames of a broad international collaboration supported by projects of INTAS, NATO Science for Peace and 6 EC Framework Program aimed at design of nanocomposite active components of catalysts for steam reforming of methane and biofuels into syngas both for the fuel cell application and synfuels production mainly published in last 5 years (Pavlova et al., 2007; Mezentseva et al., 2010; Sadykov et al., 2006c; 2008a,b; 2009a,b,c; 2010a,c; Yaseneva et al., 2008; 29-39). A lot of attention was paid to systematic studies of the effect of chemical composition of doped ceria-zirconia solid solutions on the real structure, oxygen mobility and reactivity in nanocomposites considered to be the most important factors controlling their performance. Information on the basic structural features and oxygen mobility/reactivity in Pt-supported ceria-zirconia solid solutions as well as their catalytic properties in transformation of methane and oxygenates into syngas studied earlier in details (Frolova et al., 2006; Sadovskaya et al, 2007; Sadykov et al., 2006b, 2007a-e, 2009a,d, 2010c) provide required bases for understanding the catalytic properties of nanocomposites containing these constituents.

Nanocomposites possessing promising performance and coking stability in target reactions were supported on anode substrates or heat-conducting metal substrates and successfully tested in realistic conditions in the in-cell methane steam reforming as well as in pilot-scale reactors of CH₄ or oxygenates transformation into syngas.

2. Real structure of (nano)composite materials: Effect of method of preparation on morphology and interaction between phases of composite

To provide compatibility of (nano)composites supported as porous layers on traditional Ni/YSZ cermet anodes (high ionic and electronic conductivity, close values of thermal and chemical expansion coefficients are required), the content of NiO and YSZ in composites is to be rather high. For nanocomposites supported as porous layers on heat-conducting substrates (Crofer interconnects, Ni-Al compressed foam substrates, Fechraloy foils/gauzes covered by protective corundum layer (Sadykov et al., 2008a, 2009c; 2010a), these restrictions are less severe, and the content of NiO and YSZ could be broadly varied as dependent upon the target application.

Hence, next types of (nano)composites were prepared and studied within this research program:

1. Composite I (specific surface area 11 m²/g) comprised of 60 wt.% NiO+40 wt.% YSZ was prepared by mixing and ball milling of industrial sources followed by calcination at 900 °C for 2 h (Sadykov et al., 2008b). This composite containing rather big particles of NiO and YSZ was promoted by supporting 10 wt.% of fluorite-like (Ce₀.₅Zr₀.₅O₂₋ₓ, www.intechopen.com
Pr$_{0.3}$Ce$_{0.35}$Zr$_{0.35}$O$_2$, La$_{0.3}$Ce$_{0.35}$Zr$_{0.35}$O$_2$, or perovskite-like (La$_{0.8}$Pr$_{0.2}$Mn$_{0.2}$Cr$_{0.8}$O$_3$) oxides by impregnation with water solutions of corresponding polyester citric-acid-ethylene glycol precursors followed by drying and calcination in air at 700°C for 4 h.

2. Composite II (specific surface area 23 m$^2$/g) comprised of 10 wt.% La$_{0.8}$Pr$_{0.2}$Mn$_{0.2}$Cr$_{0.8}$O$_3$+55 wt.% NiO+35 wt.% ScCeSZ was prepared using powdered Sc$_{0.1}$Ce$_{0.01}$Zr$_{0.89}$O$_{2-y}$ electrolyte synthesized by co-precipitation as described elsewhere (Smirnova et al., 2007). ScCeSZ powder was first dispersed in the water solution of Ni nitrate and polymeric electrolyte composed of perovskite-like (La$_{0.8}$Pr$_{0.2}$Mn$_{0.2}$Cr$_{0.8}$O$_3$) oxides by impregnation with complex nitrate solutions followed by drying and calcination at 800°C (Sadykov et al., 2009b).

3. Composite III (specific surface area 9 m$^2$/g) comprised of 60 wt.% NiO+40 wt.% YSZ was prepared by impregnation of powdered Y$_{0.08}$Zr$_{0.92}$O$_{2-y}$ (Russian source) with an excess of Ni nitrate solution followed by drying overnight in air at 90°C with subsequent calcination at 800°C. After regrinding, the composite III was loaded with 10 wt.% fluoro-like oxides (Pr$_x$Ce$_{3-y}$Zr$_y$O$_2$, La$_x$Pr$_{0.1}$Ce$_{3-y}$Zr$_y$O$_2$, Sm$_x$Pr$_{0.1}$Ce$_{3-y}$Zr$_y$O$_2$, where y = 0.35, 0.3, 0.2; x = 0.15-0.3; q = 0.15) by impregnation with respective mixed nitrates solutions followed by drying and calcination at 800°C for 4 h.

4. Composite IV (specific surface area 28 m$^2$/g) comprised of 10 wt.% Pr$_{0.15}$La$_{0.15}$Ce$_{0.35}$Zr$_{0.35}$O$_2$+55 wt.% NiO+35 wt.% YSZ (Russian source) was prepared by the one-pot Pechini procedure similar to that used for preparation of composite II.

5. Composites of series V comprised of 10-80% La$_{0.8}$Pr$_{0.2}$Mn$_{0.2}$Cr$_{0.8}$O$_3$+ 90-20% (NiO + YSZ) were prepared by the one-pot Pechini procedure similar to that used for preparation of composite II [38].

6. Composites of series VI comprised of 10-80% Sm$_{0.15}$Pr$_{0.15}$Ce$_{0.35}$Zr$_{0.35}$O$_2$ + 90-20% (NiO + YSZ) were prepared by the one-pot Pechini procedure similar to that used for preparation of composite II.

7. Composites of series VII comprised of 80%Pr$_{0.30-x}$Sm$_{0.70}$Ce$_{0.35}$Zr$_{0.35}$O$_2$ (x=0, 0.15, 0.3) +10%NiO + 10% Y$_{0.08}$Zr$_{0.92}$O$_{2-y}$ were prepared by the one-pot Pechini procedure similar to that used for preparation of composite II.

Pd, Pt, Ru or Pt + Ru (0.3-1.4 wt.%) were supported on composites by the incipient wetness impregnation with PdCl$_2$, H$_2$PtCl$_6$, or RuCl$_3$ solutions followed by drying and calcinations at 800°C for 2 h.

2.1 Microstructural features of (nano)composites by TEM with EDX

2.1.1 Composites with a high content of NiO and YSZ

The initial composite I is comprised of bulky NiO particles and 8YSZ aggregates with a rather good crystallinity without any apparent interaction between these oxide phases (Fig. 1). The perovskite and fluoro phases present on the NiO surface generate a moiré pattern caused by overlapping of complex oxides and NiO lattices (Fig. 2). This suggests rather good epitaxy between these phases due to a strong chemical interaction.

Composite II (Fig. 3) is comprised of loose micron-size aggregates of nanoparticles with pronounced spatial variation of their composition as revealed by EDX. In this system, particles of NiO and perovskite-like phase possessing rather good crystallinity are stacked nearly coherently. In Fig. 4, the interphase boundary a-c corresponds to the ideally matched (100) LnCrMnO$_3$ and (111) NiO planes, while the boundary b - c corresponds to stacking of (110) LnCrMnO$_3$ and (002) NiO planes with the angle of distortion equal to 5.38°. A perovskite-like phase is also present as rather disordered regions situated between the NiO particles (Fig. 5).
Fig. 1. Typical image of NiO/8YSZ composite I particles with EDX data.

Fig. 2. Typical moire patterns due to perovskite (left) of fluorite (right) layers on NiO particles in composite I.

Fig. 3. Typical TEM image of composite III aggregates with EDX data for regions 1 and 2.
Composite III is comprised of big (~1 mkm) porous aggregates formed by stacking of particles with typical sizes in the range of 10–100 nm (Fig. 6a). YSZ particles are comprised of well-crystallized coherently stacked domains with distances 3.02Å˚ corresponding to the (111) planes of cubic zirconia [JCPDS 70-4431] separated by nanopores with walls oriented along the lattice planes (Fig. 6b). In other regions (Fig. 6c) juxtaposed NiO and fluorite-like oxide particles containing elements corresponding to cations of doped ceria–zirconia oxide are revealed. A high-resolution image of some regions (Fig. 6d) shows nearly coherently intergrown particles of YSZ and NiO. EDX analysis revealed considerable incorporation of components corresponding to doped ceria–zirconia phase into YSZ, which is also reflected in the increase of (111) spacing to 3.08Å˚. Simultaneously, presence of some Ni in EDX spectrum suggests its presence in the surface layer of YSZ, perhaps, as perovskite-like clusters, though its incorporation into the lattice or nanopores of YSZ is possible as well. Traces of Zr and Y were also revealed in the regions of NiO particles.
2.1.2 Microstructure of composites with a low content of NiO and YSZ

For these composites prepared via Pechini route, big NiO particles are usually not observed (Fig. 7). Hence, in this case Ni cations appear to be mainly incorporated into the surface layer of the main phase –doped ceria-zirconia or mixed chromate-manganite. There is some non-uniformity in phases/elements distribution in nanocomposites as revealed by EDX. Broad variation of (111) spacing in fluorite-like oxide –from 3.08 to 3.13 Å is apparently caused by variation of the chemical composition of neighboring domains.

For composites with a high specific surface area supported precious metals were usually not observed by TEM as separate particles due to their high dispersion and strong interaction with complex oxides.

Hence, TEM and EDX studies revealed pronounced interaction between phases present in composites reflected in redistribution of elements between constituting phases including surface decoration and incorporation into the surface layers and in the bulk of particles. After testing in the reaction of \( \text{CH}_4 \) steam reforming in stoichiometric steam/methane feeds, NiO particles are transformed into Ni\(^\circ\). However, majority of particles remain in a close contact with the particles of complex oxide promoters. In this case their surface remains to be free of carbon. Graphite-like deposits were observed only on the surface of Ni particles not covered by complex oxide promoters (Sadykov et al., 2008b).
Fig. 7. TEM images and EDX spectra for nanocomposite 1.2%Pt/80%SmPrCeZr+20Ni/YSZ

2.2 X-ray diffraction data
For composites I and II with a high content of NiO and doped zirconia, strong reflections of these phases and weak broad reflections corresponding to dispersed complex oxide promoters were observed (Sadykov et al., 2008b, 2009b). After testing in the reaction of CH₄ steam reforming in stoichiometric feeds, NiO is completely reduced to metallic Ni. For composite II without supported Pt group metals, a weak broad reflection at 2θ ~ 27° corresponding to graphitized carbon was observed agreeing with TEM data (Sadykov et al., 2008b, 2009b). More subtle details were revealed in diffraction patterns of composite III and composites with a low content of NiO and YSZ. Thus, for composite III both before and after supporting doped ceria-zirconia oxide, along with reflections corresponding to cubic/tetragonal YSZ phase [JCPDS 70-4431] (highly dispersed c- or t-ZrO₂ phases are not discerned easily by routine XRD analysis (Sadykov et al., 2009b), reflections corresponding to monoclinic zirconia phase [JCPDS 78-1807] were observed as well (Fig. 8). As judged by the ratio of reflections intensity, promotion of NiO + YSZ composite by Pr-La-doped ceria-zirconia oxide increases further the content of monoclinic zirconia. This implies that preparation procedure of composite III, namely, successive impregnation of YSZ powder with the excess of acidic nitrate solutions followed by evaporation to dryness and calcination (Sadykov et al., 2009b) favors leaching of Y from doped zirconia. This results in destabilization and disordering of the cubic zirconia structure, while at least the surface layers of perovskite-like yttrium nickelates can be formed. Such variation of the structural features of doped zirconia was not observed for NiO + YSZ (ScCeSZ) composites promoted by complex oxides via impregnation with solutions of polymeric polyester precursor (Pechini route) even for samples with a low content of highly dispersed YSZ (Fig. 9). Hence, acidity of impregnation solutions and the time of their contact with YSZ during evaporation appear to play the main role in leaching of Y from YSZ.

Within all uncertainty of estimation of position for rather broad reflections of dispersed doped ceria-zirconia oxide, it remained more or less the same in nanocomposites as in pure oxides (Fig. 9), so redistribution of elements between YSZ and doped ceria-zirconia particles (if any) can be limited to the interfaces. The broadening of doped ceria-zirconia diffraction peaks in nanocomposite and the intensity decline suggest though some disordering of this phase due to the effect of new incoherent interfaces with NiO and YSZ. Note that reflections of NiO phase are very narrow despite its relatively low amount. Hence, at least for NiO particles observed by XRD, their size distribution is to be rather narrow as well.
Fig. 8. XRD patterns of $\text{Pr}_{0.15}\text{La}_{0.15}\text{Ce}_{0.35}\text{Zr}_{0.35}\text{O}_2$ (1), 10% $\text{Pr}_{0.15}\text{La}_{0.15}\text{Ce}_{0.35}\text{Zr}_{0.35}\text{O}_2$ / NiO + YSZ (2) and NiO + YSZ (3). ● Ce$_{0.35}$Zr$_{0.35}$La$_{0.15}$Pr$_{0.15}$O$_2$, | - Zr(Y)O$_2$ monoclinic, ▼ - Zr(Y)O$_2$ cubic, ▼ - NiO.

Fig. 9. XRD patterns of Zr$_{0.92}$Y$_{0.08}$O$_2$ calcined at 500 °C (1), Sm$_{0.15}$Pr$_{0.15}$Ce$_{0.35}$Zr$_{0.35}$O$_2$ complex oxide (2) and composite 80% Sm$_{0.15}$Pr$_{0.15}$Ce$_{0.35}$Zr$_{0.35}$O$_2$/10%NiO/10% Zr$_{0.92}$Y$_{0.08}$O$_2$ (3). *- Zr$_{0.92}$Y$_{0.08}$O$_2$, +-Sm$_{0.15}$Pr$_{0.15}$Ce$_{0.35}$Zr$_{0.35}$O$_2$.

For Pt-promoted composites with a low content of NiO, rather narrow reflections corresponding to Pt particles were detected (Fig. 10). As was earlier shown for Pt-supported doped ceria-zirconia nanocrystalline oxides (Sadykov et al., 2007e), only a small part of supported Pt is present as metal particles detected by XRD, while oxidic forms (clusters, Pt$^{2+}$ cations) stabilized due to strong interaction with support dominate. Note that supported Ru is not detected by XRD (Fig. 10) due to a higher stability of its oxidic forms.

Hence, structural studies of nanocomposites comprised of NiO, YSZ, complex oxide promoters and supported Pt group metals revealed pronounced interaction between constituting phases. Decreasing the sizes of constituting phases and modification of preparation procedure favor this interaction manifested as heteroepitaxy, decoration and redistribution of elements between neighboring domains. One-pot Pechini procedure provides the smallest sizes of constituting phases particles, while impregnation with acidic nitrate solutions favor redistribution of elements between phases due to leaching.
3. Reactivity of (nano)composites

3.1 H$_2$ TPR

Reduction of NiO in composites by H$_2$ proceeds via the topochemical mechanism, which includes generation of Ni$^+$ nuclei at some specific surface sites (outlets of dislocations etc) with their subsequent growth accompanied by the expansion of reaction area at the NiO/Ni$^+$ interface (and, respectively, reaction rate increase) until nuclei overlap forming a dense layer of Ni$^+$ product followed then by the reaction rate decline described by a contracting sphere model (Boldyrev et al., 1979). Hence, this reaction is sensitive to the real/defect structure, dispersion and surface composition of NiO particles in green composites, which are of a great importance for understanding the catalytic properties of these systems in steam reforming of fuels. Oxide additives should not cover all the surface of NiO particles by a dense layer to cause noticeable effect on their reduction characteristics. It is sufficient to decorate a small number of surface defect sites in vicinity of the dislocation outlet or domain/grain boundaries to hamper or accelerate removal of oxygen from these sites, and, hence, dynamics of Ni$^+$ nucleation, thus shifting TPR peak position.

Typical H$_2$ TPR spectra for NiO-doped zirconia nanocomposites, both initial and with supported complex oxides, are shown in Figs. 11, 12. For all samples, NiO reduction was completed up to 600–700 °C. For composite I, reduction starts at ~280 °C with $T_{\text{max}}$ ~ 350 °C, which is rather close to typical characteristics for samples of pure NiO or its mechanical mixture with YSZ (Montoya et al., 2000; Sanchez-Sanchez et al., 2007) and agrees with the microstructural data revealing little if any interaction between NiO and YSZ in this case. Supporting Ce–Zr–O oxide on this composite shifts $T_{\text{max}}$ to ~400 °C, apparently due to a partial blocking of NiO surface (especially surface defects at which nucleation of Ni$^+$ occurs) and/or stabilizing Ni$^{2+}$ cations thus hampering reduction. Indeed, similar position of reduction peak was earlier observed for NiO supported on Ce-Zr-O$_2$ (Romero-Sarria et al, 2008).
Fig. 11. Typical $\text{H}_2$ TPR spectra for different NiO/YSZ composites promoted by fluorite-like oxides (left) or perovskite-like oxides (right). 5 % $\text{H}_2$ in Ar, 5 l/h, temperature ramp 5ºC/min. Left: 10% $\text{Ce}_0.5\text{Zr}_0.5\text{O}_2$/composite I (1), composite I (2), composite VI (3), 10% SmPrCeZr/composite III (4). Right: 0.3% Pd/10% LaPrMnCr/composite I (1), composite II (2), 0.3% Pd/composite II (3).

For unpromoted NiO + YSZ composite III, reduction starts at $\sim$330 ºC and is characterized by three overlapping peaks with $T_{\text{max}}$ situated at $\sim$ 350, 400 and 450 ºC (Sadykov et al., 2009b). In addition, the high-temperature (up to 700ºC) tail of hydrogen consumption was observed for both undoped and doped samples of composite III. The first peak can be assigned to reduction of NiO particles not modified by interaction with YSZ. The second peak is close by position to that in composite I promoted by $\text{Ce}_0.5\text{Zr}_0.5\text{O}_2$ oxide (Fig. 11), so it can be similarly explained by decoration of the surface of NiO particles with irreducible $\text{Y}($Zr$)\text{O}_x$ oxidic species. Indeed, TPR peaks with $T_{\text{max}}$ situated even at higher (450–600ºC) temperatures were observed in the case of NiO supported on zirconia, ceria–zirconia or alumina (Montoya et al., 2000; Srinivas et al., 2003; Matsumura & Nakamori, 2004). By analogy with these data, the high-temperature tail of hydrogen consumption can be assigned to reduction of Ni cations incorporated into the bulk of doped zirconia particles (vide supra). Hence, TPR data revealed pronounced variation of the reactivity of NiO particles/species even in unpromoted NiO + YSZ composites, which could reflect different degree of chemical interaction between components.

Supporting doped ceria-zirconia complex oxides on composite III practically eliminates the first TPR peak situated at $\sim$350ºC (Fig. 11), slightly shifts the second peak to higher temperatures and increases the relative share of the high-temperature peak at $\sim$470ºC. A high-temperature reduction tail remains to be observed for all samples. This implies that the second impregnation provides modification of the surface of all NiO particles present in composite by Zr and rare-earth cations.

For composite VI prepared via Pechini route with the same type of PrSmCeZrO complex oxide additive and having much smaller content of NiO and YSZ only one $\text{H}_2$ TPR peak (Fig. 11, curve 3) was observed. A lower intensity of this peak as compared with that for other samples (curves 1 and 4, Fig. 11) correlates with a smaller NiO content, while its position is rather close to that for other peaks assigned to reduction of NiO particles decorated by fluorite-like oxides. This suggests that reactivity and lattice oxygen mobility of small NiO particles strongly interacting with complex fluorite-like is rather close to that of bulky particles decorated with fragments of fluorite-like oxides.
Complex perovskite oxide strongly hampers NiO reduction in the composite I (Fig. 11, curve 1): reduction peak shifts to \( \approx 460^\circ C \) even despite co-promotion with Pd. This is explained by the relatively low reducibility/lattice oxygen mobility of complex manganochromite (Peña-Martínez et al., 2006; Sfeir et al., 2001; Sauvet & Irvine, 2004) rather uniformly covering the surface of NiO particles in this composite thus hampering reduction. For composite II also containing complex perovskite phase as an additive and prepared via one-pot Pechini route, the reduction peak is shifted to even higher (\( \approx 530^\circ C \)) temperatures (Fig. 11, curve 2). In agreement with TEM data (vide supra) this suggests even stronger interaction between the perovskite and NiO particles in this sample. Appearance of the high-temperature shoulder at \( \approx 600^\circ C \) implies that some Ni cations could be even incorporated into the bulk of perovskite particles/domains thus forming Ni-substituted oxide (Sauvet & Irvine, 2004; Sfeir et al., 2001). Promotion of this composite with a small amount of Pd shifts reduction peak to lower temperatures and removes high-temperature shoulder (Fig. 11, curve 3).

In general, promoting effect of supported Pt group metals on the oxides reduction by hydrogen is a well-documented phenomenon explained by the efficient activation of \( H_2 \) molecules on the metal particles and spill-over of atomic hydrogen onto the oxide surface, thus easily removing reactive oxygen forms (Bernal et al., 2002). Downward shift of \( H_2 \) TPR peak is observed for any mechanism of the solid oxide reduction, either topochemical (such as NiO reduction) or diffusion-controlled (reduction of ceria-zirconia solid oxide solution). Facilitation of the lattice oxygen mobility due to incorporation of precious metals into domain boundaries and subsurface layers is demonstrated as well (Sadovskaya et al., 2007).

Promotion of composites by Ru or Pt clearly accelerates their bulk reduction by hydrogen (Figs. 12) characterized by the main peak situated at 370–400\(^\circ C\). In addition, in the low-temperature (100–300\(^\circ C\)) range, new peaks appear apparently corresponding to reduction of different Ru and/or Pt oxidic species accompanied by reduction of complex fluorite-like oxides. For Ru supported on ceria or zirconia, peaks assigned to reduction of RuO\(_2\) species with different dispersion are situated at 80–100\(^\circ C\) [Hosokawa et al., 2003; Yan et al., 2007]. For Pt-supported Pr-Ce-Zr-O samples, reduction starts at \( \approx 150^\circ C \) with the main maximum situated at \( \approx 280^\circ C \) and a shoulder at \( \approx 200^\circ C \) (Sadykov et al., 2007e). Hence, TPR peaks at 115–150\(^\circ C\) observed for Ru-promoted samples of composites with a high NiO/YSZ content (Sadykov et al., 2009b) suggest rather weak interaction of RuOx species with other phases present in composites. When Ru cations are incorporated into the surface vacancies of La–Sr–chromate with the perovskite structure, these peaks are shifted to higher (180–200\(^\circ C\)) temperatures (Yan et al., 2007). Hence, peak observed at \( \approx 200^\circ C \) for nanocomposite with a high (80\%) content of Pr-Ce-Zr-O fluorite can be assigned to reduction of RuO\(_x\) species strongly interacting with this oxide, perhaps, modified also by dissolved Ni cations (Fig. 12).

This interaction apparently facilitates reduction of NiO present in promoted composites, since \( T_{\text{max}} \) of the main reduction peak is shifted downward. For Pt-supported nanocomposite (Fig. 12), position and shape of the low-temperature peak is practically the same as that observed for Pt/Pr-Ce-Zr-O samples (Sadykov et al., 2007e). Intermediate position is observed for Pt+Ru-promoted samples (Fig. 12) suggesting formation of mixed PtRuOx oxidic species.
3.1 TPR by fuel molecules (CH$_4$, C$_2$H$_5$OH) and reoxidation by H$_2$O
Temperature-programmed reduction of composites by fuels (methane, ethanol) followed by reoxidation with H$_2$O characterizes mobility and reactivity of oxygen species and provides also an additional information on ability of surface sites to activate reagents.

3.1.1 CH$_4$ TPR
Reduction of bulk NiO sample (a starting compound for preparation of composite I) starts at ~400°C and occurs in a narrow (~100°C) temperature range with practically simultaneous appearance of deep (CO$_2$, H$_2$O) and partial (CO, H$_2$) oxidation products (Fig. 13). While CO$_2$ evolution rapidly falls to zero, other products are observed up to 800°C. H$_2$ evolution declines to zero when sample is completely reduced. This means that bulk Ni$^0$ particles are not able to continuously dissociate CH$_4$, apparently due to a rapid surface blocking by the graphitic carbon.

Fig. 13. Typical CH$_4$ TPR data for NiO (left) and nanocomposite III (right). Experimental parameters for Figs. 13-14: contact time 0.1 s, 1% CH$_4$ in He, temperature ramp 5°C/min.
For undoped composites, reduction by CH$_4$ starts at higher than for pure NiO temperatures (Fig. 13), which agrees with H$_2$ TPR data (vide supra) and is explained by migration of irreducible cations from YSZ onto the surface of NiO. Some H$_2$ evolution continues even after complete reduction of composites due to CH$_4$ decomposition. This suggests that interaction between components even in undoped composites prevents formation of a dense layer of graphitic carbon.

Doping by complex oxides and Pt group metals facilitates the reduction shifting TPR peaks to lower temperatures (Fig. 14). This is explained by a higher efficiency of Pt group metals in C-H bond dissociation as compared with the centers of NiO or fluorite-like/perovskite-like oxides. Indeed, reduction of doped ceria or ceria–zirconia oxides by methane starts from ~400°C with evolution of deep oxidation products followed by syngas generation at higher (>700°C) temperatures (Sadykov et al., 2007a). As far as CH$_4$ pyrolysis is concerned, there appears to be some synergy between the action of Pt group metal, complex oxide additive and Ni. Hence, interaction between components in nanocomposite apparently helps to prevent formation of dense graphitic layers in the course of CH$_4$ pyrolysis. Polymerized C$_x$H$_y$O$_z$ species inevitably formed on the surface of nanocomposites due to this pyrolysis appear to possess more loose structure due to incorporation of oxygen atoms. Reoxidation of deposited coke by H$_2$O revealed that for Pt-supported doped ceria–zirconia sample CO$_2$ evolution in the course of H$_2$O oxidation starts at temperatures below 400°C without any evolution of H$_2$ up to ~ 550°C (Fig. 15). It clearly can be explained only by decomposition of some surface carbonate complexes - either organic ones within coke precursors or inorganic ones. The maximum of CO, CO$_2$ and H$_2$ evolution due to oxidation of coke by water accompanied by simultaneous reoxidation of doped ceria–zirconia is situated at ~ 880°C (Fig. 15). For Ni/YSZ composites, both promoted or not, oxidation of deposited during CH$_4$ TPR runs carbonaceous species accompanied by simultaneous CO$_2$, CO and H$_2$ evolution starts earlier and proceeds faster than for Pt-supported fluorite-like oxides (Fig. 15). This suggests involvement of Ni surface atoms together with Pt in gasification of the surface coke. Indeed, while Ni atoms can be oxidized by H$_2$O producing hydrogen and oxygen atoms, it is impossible for Pt atoms. So, the function of Ni in this case is to provide efficient activation of mild oxidant- water at temperatures lower than those typical for reduced doped ceria-zirconia oxide. On the other hand, Pt atoms could use these oxygen atoms supplied to them via spillover for the efficient oxidation of coke precursors.

Fig. 14. Typical CH$_4$ TPR data for samples of 1.4Pt/10Pr$_{0.15}$Sm$_{0.15}$Ce$_{0.5}$Zr$_{0.2}$O$_2$/composite III (left) and 1Ru/10LaMnCrPr/composite V.
Another new feature revealed in these experiments is evolution of CH\textsubscript{x} (or, in general, hydrocarbons) in the process of H\textsubscript{2}O TPO of reduced and coked promoted composite (Fig. 15). This supports hypothesis that complex oxide promoters make deposited coke species less dense and, perhaps, containing a bigger fraction of hydrogen atoms (Sadykov et al., 2008b), so they are more easily cracked by interaction with the surface hydroxyls. In H\textsubscript{2}O TPO spectra of both promoted and unpromoted composites (Fig. 15) peaks of CO\textsubscript{2} evolution in the isothermal mode at 880ºC are present. A similar by position peak is present in the H\textsubscript{2}O TPO spectrum of Pt-supported coked doped ceria-zirconia oxides (Fig. 15, left). Hence, its appearance in the case of nanocomposites TPO spectra can be explained by oxidation of a part of carbonaceous species located on the surface of YSZ or doped ceria-zirconia particles remote from the interface with Ni particles.

### 3.1.2 C\textsubscript{2}H\textsubscript{5}OH TPR

In these experiments, to prevent condensation of unreacted ethanol and its partial oxidation products in the gas analyzer, a cooled to -40ºC trap was situated at the reactor exit. This precludes estimation of carbon balance and degree of ethanol conversion. However, analysis of dynamics of products evolution allows to make useful conclusions about reactivity and route of transformation of ethanol for different types of composites.

Figs. 15, 16 show typical results of ethanol TPR experiments for composites with broadly varying compositions. In general, their reactivity with respect to ethanol is comparable: at ~300ºC H\textsubscript{2} appears followed by other products including H\textsubscript{2}O, CO\textsubscript{2}, CO and CH\textsubscript{x}. While CO\textsubscript{2} and H\textsubscript{2}O are products of ethanol oxidation by the lattice oxygen of composites, CH\textsubscript{x} is comprised of non-condensed products passed through the trap such as CH\textsubscript{4} and C\textsubscript{2}H\textsubscript{4} formed due to cracking and dehydration of ethanol (de Lima et al., 2008). After complete reduction of catalysts, evolution of H\textsubscript{2} and CO continues due to decomposition of ethanol (Domine et al., 2008). Though it is inevitably accompanied by accumulation of surface carbonaceous species, however, within the time scale of experiments, performance is stable which suggests deposition of loose coke species. Since in all cases at the steady state the H\textsubscript{2}/CO ratio in the products is close to 2.5, by stoichiometry of ethanol decomposition this coke should retain a lot of hydrogen atoms. CH\textsubscript{x} formation due to ethanol cracking continues only in the case of composites containing doped ceria-zirconia oxides (Fig. 15, left). In studied conditions concentrations of CO (~0.5-0.75%) and H\textsubscript{2} (~1.25-1.75%) were rather close for studied composites.

Temperature-programmed reoxidation by H\textsubscript{2}O of deposited carbonaceous species starts at rather low (~400ºC) temperatures and proceeds fast (Fig. 16). Hence, for nanocomposites containing even relatively low content of complex oxide promoters carbonaceous deposits derived from ethanol are highly reactive, which is one of the criteria of stability of these composites performance in steam reforming of ethanol (de Lima et al., 2008).

### 4. Catalytic activity in SR of CH\textsubscript{4} and biofuels

#### 4.1 Catalytic properties of dispersed nanocomposites in methane steam reforming

#### 4.1.1 Composites with fluorite-like oxide additives.

Typical values of CH\textsubscript{4} conversions for different nanocomposites and estimated from these values effective first-order rate constants are given in Figs. 17, 18 and Table 1.
Fig. 15a. Temperature-programmed oxidation by H\textsubscript{2}O of reduced and coked in CH\textsubscript{4} TPR runs samples of 1.4 wt.% Pt/Pr\textsubscript{0.15}Sm\textsubscript{0.15}Ce\textsubscript{0.5}Zr\textsubscript{0.2}O\textsubscript{2} (left) and 1.4 wt.% Pt/Pr\textsubscript{0.15}Sm\textsubscript{0.15}Ce\textsubscript{0.5}Zr\textsubscript{0.2}O\textsubscript{2}/composite III (right) samples. 1% H\textsubscript{2}O in He, 5\textdegree/min, 0.1 s.

Fig. 15b. Typical EtOH TPR spectrum for sample of nanocomposite series VI promoted with 80%SmPrCeZr (left) and nanocomposite series V promoted by 80%LaPrMnCr (right). Experimental conditions for Figs. 15-16: 1% EtOH in He, contact time 0.1 s, temperature ramp 5\textdegree/min.

Fig. 16. EtOH TPR (left) followed by H\textsubscript{2}O TPO (right) for nanocomposite series V promoted by 1%Ru and 10%LaPrMnCr.
Fig. 17. Temperature dependence of CH\textsubscript{4} conversion (left) and effective first-order rate constants (right) for different composites at 10 ms contact time. Left: 1-10 wt.% Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite \textit{III}, 2-1.3 wt.% Ru/10 wt.% Pr\textsubscript{0.15}La\textsubscript{0.15}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite \textit{III}, 3-0.3 wt.% Pt/10 wt.% Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite \textit{I}, 4- composite \textit{IV}. Feed 8% CH\textsubscript{4}+ 8% H\textsubscript{2}O in Ar. Right: 1-composite \textit{I}, 2-Ce\textsubscript{0.5}Zr\textsubscript{0.5}O\textsubscript{2}/composite \textit{I}, 3-0.3%Pd/10%Ce\textsubscript{0.5}Zr\textsubscript{0.5}O\textsubscript{2}/composite \textit{I}, 4,5-0.3%Pt/10% Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite \textit{I}. Feed 8% CH\textsubscript{4}+ 8% H\textsubscript{2}O (1-4) or 8% CH\textsubscript{4}+ 24%H\textsubscript{2}O (5).

Supporting complex ceria-zirconia oxides increases catalysts performance at temperatures exceeding 600ºC and stabilizes it at high temperatures (Figs. 17, Table 1) apparently due to hampering of coking. Some samples based upon fluorite oxides promoted composite \textit{III} remain inactive at 650ºC (Table 1), while composite \textit{IV} demonstrates a good performance starting already from 550ºC (Fig. 17). This can be explained both by a higher specific surface area of composite \textit{IV} as well as by a higher degree of interaction between components of this composite provided by one-pot Pechini route of synthesis, and, hence, stabilization of small active Ni clusters on the surface of oxidic components of composite (vide supra).

For the stoichiometric feed, Pd as co-promoter further increases performance of ceria-zirconia promoted composite \textit{I} (Fig. 17, Table 1) making it highly active even at temperatures below 600ºC. Pt as co-promoter also ensures a high activity in the middle-temperature range (Fig. 17, Table 1). Indeed, at 650ºC, effective first-order rate constants estimated for the plug-flow reactor are as high as 30-40 s\textsuperscript{-1} (Fig. 17). Hence, more efficient activation of CH\textsubscript{4} molecules on Pt or Pd clusters demonstrated by CH\textsubscript{4} TPR experiments allows to increase the overall performance of Ni-containing catalysts in the intermediate temperature range, while dispersed fluorite-like oxide promoter along with Ni apparently plays the major role in activation of water molecules and transfer of activated oxygen-containing species (hydroxyls, hydroxocarbonates) to the sites where activated CH\textsubscript{x} species are located, thus preventing formation of coke.

The performance of samples co-promoted with Pt and fluorite-like oxides tends to increase in feeds with the excess of steam (Figs. 17, Table 1). This is important from the practical point of view since in real SOFC operation conditions, the oxygen ions transfer through the cell increases the overall content of oxidants within the porous composite anode. The increase of steam content in the feed increases the rate of activated hydrocarbons species transformation into syngas, thus preventing the surface coking and ensuring a high performance of composite co-promoted with Pt and fluorite-like oxide. For Pt as a co-promoter, the effect of the steam excess in feed strongly depends on the exact chemical
## Table 1. Values of CH\textsubscript{4} conversion for CH\textsubscript{4} SR at 650°C on composites promoted with fluorite/perovskite-like oxides and Pt, Pd or Ru. 10 ms contact time, 1:1 feed (8% CH\textsubscript{4} + 8% H\textsubscript{2}O in He), 1:3 feed (8% CH\textsubscript{4} + 24% H\textsubscript{2}O in He) and 1:2 feed* (20% CH\textsubscript{4} + 40% H\textsubscript{2}O, Ar balance).

<table>
<thead>
<tr>
<th>Composition</th>
<th>CH\textsubscript{4} conversion, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH\textsubscript{4}: H\textsubscript{2}O</td>
<td>1:1</td>
</tr>
<tr>
<td>Ce\textsubscript{0.5}Zr\textsubscript{0.5}O\textsubscript{2}/composite I</td>
<td>23</td>
</tr>
<tr>
<td>1.4 wt% Pt/La\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}</td>
<td>15</td>
</tr>
<tr>
<td>0.3 wt% Pd/Ce\textsubscript{0.3}Zr\textsubscript{0.5}O\textsubscript{2}/composite I</td>
<td>43</td>
</tr>
<tr>
<td>0.3 wt% Pt/La\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite I</td>
<td>30</td>
</tr>
<tr>
<td>0.3 wt% Pt/Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite I</td>
<td>35</td>
</tr>
<tr>
<td>1.03 wt% Pt/Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite I</td>
<td>0</td>
</tr>
<tr>
<td>0.5 wt% Ru/Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite I</td>
<td>12</td>
</tr>
<tr>
<td>0.3 wt% Pd/LaMnCrPr/composite I</td>
<td>26</td>
</tr>
<tr>
<td>1 wt% Ru/LaMnCrPr/composite I</td>
<td>70</td>
</tr>
<tr>
<td>Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite III</td>
<td>0</td>
</tr>
<tr>
<td>1 wt% Ru/Pr\textsubscript{0.3}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite III</td>
<td>32</td>
</tr>
<tr>
<td>1 wt% Ru/Pr\textsubscript{0.15}La\textsubscript{0.15}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite III</td>
<td>36</td>
</tr>
<tr>
<td>1 wt% Ru/Pr\textsubscript{0.15}Sm\textsubscript{0.15}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}/composite III</td>
<td>19</td>
</tr>
<tr>
<td>Composite IV (10Pt\textsubscript{0.15}La\textsubscript{0.15}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}+55NiO+35YSZ)</td>
<td>40</td>
</tr>
<tr>
<td>Composite V\textsubscript{1} 10LaPrMnCr+90Ni/YSZ (4.1 m\textsuperscript{2}/g)</td>
<td>21</td>
</tr>
<tr>
<td>Composite V\textsubscript{2} 50LaPrMnCr+50Ni/YSZ (8.1 m\textsuperscript{2}/g)</td>
<td>20</td>
</tr>
<tr>
<td>Composite V\textsubscript{3} 80LaPrMnCr+20Ni/YSZ (12 m\textsuperscript{2}/g)</td>
<td>22</td>
</tr>
<tr>
<td>1%Ru/composite V\textsubscript{1} (3 m\textsuperscript{2}/g)</td>
<td>70</td>
</tr>
<tr>
<td>1%Ru/composite V\textsubscript{2} (8.9 m\textsuperscript{2}/g)</td>
<td>45</td>
</tr>
<tr>
<td>1%Ru/composite V\textsubscript{3} (6.2 m\textsuperscript{2}/g)</td>
<td>18</td>
</tr>
<tr>
<td>0.7 % Ru/composite V\textsubscript{2}</td>
<td>72*</td>
</tr>
</tbody>
</table>

Note also that for Pt-supported fluorite-like oxides without Ni addition performance is independent upon the water excess (Table 1). A non-additive increase of performance due to supporting Pt along with fluorite-like oxides on Ni-containing composite (a synergetic effect) is thus apparent.

For Pd supported on a composite promoted by ceria-zirconia, the increase of steam content in the feed suppresses the low-temperature performance (Table 1), which could be explained by stabilization of less reactive oxidized Pd species by fluorite-like oxide. Here the main role is played by a higher stability of Pd oxidic forms (perhaps, some surface phases including both Pd and Ce cations) as compared with those of Pt. A lower ability of oxidized Pd species to activate methane is thus responsible for a low activity in feeds with the steam excess.

For Ru as a co-promoter of composite I in combination with the fluorite-like oxides, the excess of steam also positively affects the low-temperature performance (Table 1). However, this performance is lower than for Pt-containing samples in all feeds, so Pt is more efficient co-promoter than Ru in combination with complex fluorite-like oxides. This can be explained by a higher efficiency of Pt in activation of methane (Wei & Iglesia, 2004).
For samples promoted by the same amount of Pt, activity is also higher for sample with a higher Ni content despite a lower specific surface area. Hence, the efficiency of the composite in CH\(_4\) SR is defined by the content of components (Ni, Ru, Pt) activating fuel molecules. On the other hand, when Pt or Pd are supported on Ni/YSZ composite without oxide additives, methane conversion is not improved (Sadykov et al, 2006c). This suggests that namely combination of precious metals with oxide promoters helps to provide enhanced activity of composites in the intermediate temperature range. Due to a low content of supported precious metals, their effect can be assigned to modification of some specific defect centers of Ni particles. This suggestion agrees with a high performance of composite IY prepared via one-pot Pechini synthesis procedure (Fig. 17) even without promotion by precious metals.

The synergy of the catalytic action of components in nanocomposites is retained for feeds with the excess of steam. Thus, for nanocomposites containing Pt and SmPrCeZr as co-promoters, sample with a higher Ni content provides significantly higher CH\(_4\) conversion in the whole temperature range in both stoichiometric (1:1) and more oxidizing (1:3) feeds (Fig. 18). At a low Ni content, the excess of steam provides a higher CH\(_4\) conversion in the whole temperature range due to efficient activation of CH\(_4\) on Pt sites remaining in the metallic state even in the oxidizing conditions. At a high Ni content, a complex temperature - dependent effect of the steam excess on activity suggests interplay of several factors determined by interaction between Pt, Ni and fluorite-like oxide and affecting efficiency of CH\(_4\) and H\(_2\)O activation and, hence, steady-state oxidation degree of Ni surface and its coverage by coke.

![Fig. 18. Effect of water excess in the feed on performance of nanocomposites promoted by Pt and SmPrCeZr (left) or Ru and LaPrMnCr (right) in CH\(_4\)SR and H\(_2\)O/CH\(_4\)=1 (a) or 3(b). Contact time 10 ms, CH\(_4\) concentration 8%. Left: 1-Pt/10SmPrCeZr/Ni/YSZ, 2-Pt/80SmPrCeZr/Ni/YSZ. Right: 1- Ru/50LaPrMnCr/Ni/YSZ, 2- 50LaPrMnCr/Ni/YSZ.](image)

### 4.1.2 Composites with perovskite-like oxide additives

Ni/ScCeSZ cermet without oxide promoters was not able to provide a steady-state performance in the methane SR in stoichiometric feeds due to very fast coking leading finally to the reactor plugging (Sadykov et al., 2006c, 2008b). After reduction by H\(_2\) composite II comprised of Ni/ScCeSZ cermet promoted by perovskite-like oxide demonstrates a reasonably high and stable activity at temperatures exceeding 650\(^\circ\)C (Sadykov et al., 2008b). Similarly, promotion of composite I by perovskite oxide allows to obtain a reasonable
performance in methane steam reforming even at 650 °C (Table 1). By itself, La-Pr-Mn-Cr-O perovskite is inactive in SR of methane. However, incorporation of Ni cations into the lattice of such irreducible perovskites provides a reasonable level of activity in methane steam reforming (Sauvet & Irvine, 2004). Hence, observed high and stable activity of perovskite-promoted composites in CH₄ SR in stoichiometric feed can be explained by a partial dissolution of NiO in the acidic polyester solution at the preparation stage. At subsequent calcination, Ni-containing perovskite is formed (vide supra EDX data). Under contact with reducing reaction media, small highly reactive Ni clusters are segregated on the surface of perovskite oxide being stabilized by interaction with this matrix stable in reducing conditions. In addition, the surface of big NiO particles in non-reduced composite is covered by the perovskite layers (vide supra). Hence, big Ni⁰ particles generated due to NiO reduction in the reaction media are decorated by perovskite-like oxidic species. A high efficiency of these species and/or separate perovskite particles in activation of steam and carbon dioxide facilitates gasification of CH₄ species produced by methane activation on Ni, thus preventing coking. This level of high-temperature activity for perovskite-promoted composite I (Table 1) is close to that provided by ceria-zirconia oxide promoter. Hence, similar factors affecting surface properties of Ni particles in promoted composites as well as enhanced activation of water and/or CO₂ molecules on the surface sites of oxide additives could operate in the case of both types of oxide promoters ensuring high and stable performance of composites in methane SR in stoichiometric feeds, especially at rather high (>700°C) temperatures.

The increase of steam/methane ratio from 1 to 3 was found to decrease the high-temperature performance of perovskite-promoted composite I (Sadykov et al., 2006c, 2008b). This can be assigned to partial oxidation of Ni surface atoms contacting with perovskite species/particles and their stabilization as less active perovskite-like fragments.

Co-promotion of composites with perovskite-like oxide and Ru allowed to achieve a high level of middle-temperature activity in feeds with different steam/methane ratios (Table 1, Fig. 18). For composite II, the increase of Ru content from 0.3 to 1 wt.% decreases activity in the stoichiometric feed and increases it in the feed with the excess of steam (Sadykov et al., 2010b). This behavior can be explained by a higher rate of CH₄ activation on the surface of sample with a higher Ru content, thus leading to deactivation due to coking in stoichiometric feed. In a feed with the excess of steam, enhanced rate of activated CH₄ species gasification prevents coking and provides a higher performance of sample with a higher Ru content due to a higher rate of CH₄ activation. Similar features observed for composite I co-promoted with fluorite-like oxide and Pt (Table 1) agree with this explanations of the precious metal content and feed composition effects.

For composites series V containing LaPrMnCr and promoted by 1%Ru (Table 1), activity increases with Ni content, thus clearly demonstrating synergy of Ni+ Ru action due to suppression of coking ability of Ni sites explained by the surface alloys formation. Activation of H₂O molecules on Ni atoms can be important as well.

As follows from Fig. 18, Ni/YSZ cermet promoted only by perovskite-like complex oxide provides the same CH₄ conversion in both types of feed, while co-promoting effect of Ru is much stronger in the stoichiometric feed. This suggests that in feed with the steam excess Ru is in part transformed into less active oxidic forms stabilized by perovskite-like complex oxide.

When comparing two series of samples co-promoted with precious metals and either fluorite-like or perovskite-like oxides (Table 1), the middle-temperature performance of the best samples of both series is comparable. For samples promoted with perovskite-like oxide,
the highest activity is provided by supporting Ru, while for samples with fluorite-like oxide additives, the highest performance is observed after loading Pt. Hence, these two types of systems can be considered as promising for the practical application. In general, a better performance is provided by YSZ-containing catalysts (Sadykov et al., 2010b). Though more detailed studies are required for elucidating the exact role played by the rare-earth cations in determining catalytic properties of these nanocomposites, it is clear that observed trends are not determined by the oxygen mobility in doped zirconia particles which is higher for ScCeSZ (Smirnova et al., 2007).

Hence, the most efficient promoter for Ni/YSZ-based composite catalyst is Pr(Sm)-doped ceria-zirconia in combination with Pt. Since Pt is more expensive than Ru, the latter was also used as a co-promoter in combination with either perovskite or fluorite oxide when supporting thin layers of NiO+YSZ-based nanocomposites on different substrates (vide infra). As far as the preparation procedures are concerned, one-pot Pechini route apparently provides the highest activity of promoted composites, at least, for systems with complex fluorite-like oxide additives. Hence, this method was selected as a basic one for preparation of dispersed promoted composites for subsequent supporting on different substrates. As follows from Fig. 19, performance of nanocomposite active components prepared via this route is sufficiently stable in feeds with a small excess of steam.

Temperature-programmed oxidation of nanocomposite samples discharged from reactor after testing in stoichiometric feeds demonstrated that combined action of precious metals and complex oxides decreases the amount of deposited carbon providing thus high and stable performance (Sadykov et al., 2006c, 2008b).

![Fig. 19. Stability test for nanocomposite 10%Sm_{0.15}Pr_{0.15}Ce_{0.35}Zr_{0.35}+ 55%NiO+35%YSZ +1.4%Pt at 600 °C, feed 20% CH\textsubscript{4} + 40% H\textsubscript{2}O, Ar balance, contact time 60 ms.](image)

4.2 Catalytic properties of dispersed nanocomposites in ethanol steam reforming
In this reaction, in general, in the middle-temperature range, activity of all composites, promoted by Ru or not, was rather high (Table 2, Fig. 20). This is explained by a high activity of Ni in this reaction provided coking of its surface is hindered by oxidic promoters. Moreover, specific activity of Ni supported on alumina is reported to be even higher than that of low-loaded Ru (Fatsikostas et al., 2002; Haryanto et al., 2005; Sanchez-Sanchez et al., 2007). This agrees with the results of Srinivas et al. (Srinivas et al., 2003) demonstrating a
high and stable activity in ESR of composites comprised of 40wt. % NiO and 60 wt.% CeO$_2$ (CeO$_2$-ZrO$_2$) prepared via hydrothermal route. Hence, promoted nanocomposites efficient and stable in steam reforming of methane are also quite promising for the steam reforming of ethanol. As can be inferred from the hydrogen content in converted feed, at 700 °C, even at short contact time, up to 70% of ethanol is converted along the steam reforming route. C$_2$H$_4$ and CH$_4$ were not revealed in the products, while they were inevitably present for catalysts comprised of precious metals supported on alumina (Erdőhelyi et al., 2006; Romero-Sarria et al., 2008). Hence, for developed composites containing YSZ and complex fluorite-oxide promoters, primary route of ethanol transformation into syngas apparently does not include ethanol dehydration or cracking. This agrees with very high CO$_2$/CO ratio in products at 700 °C in diluted feed clearly not conforming to the equilibrium composition of converted feed (Erdőhelyi et al., 2006; Romero-Sarria et al., 2008) or to results obtained with a high content of ethanol in the feed (vide infra). Absence or very low concentration of CO in products for catalysts comprised of precious metals supported on alumina (Erdőhelyi et al., 2006; Romero-Sarria et al., 2008) implies that red-ox properties of promoters are also responsible for routes of ethanol transformation into carbon oxides (Erdőhelyi et al., 2006; Vargas et al., 2005). Since at high temperatures acetates are the only surface species detected by infra-red spectroscopy (Erdőhelyi et al., 2006), at short contact times CO$_2$ can be considered as primary product of their transformation. Since CH$_4$ was not observed in products, this transformation is not simple cracking usually accompanied by CH$_4$ formation, but should include interaction of ethoxide species with reactive hydroxyls or hydroxocarbonate species.

<table>
<thead>
<tr>
<th>№</th>
<th>Sample</th>
<th>Products concentration. %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>CO</td>
</tr>
<tr>
<td>1</td>
<td>10% Pr$<em>{0.15}$Sm$</em>{0.15}$Ce$<em>{0.35}$Zr$</em>{0.35}$O$_2$+55% NiO+35%YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>2</td>
<td>80% Pr$<em>{0.15}$Sm$</em>{0.15}$Ce$<em>{0.35}$Zr$</em>{0.35}$O$_2$+10% NiO+10%YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>3</td>
<td>50%LaPrMnCr+30%NiO+20%YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>4</td>
<td>80%LaPrMnCr+10%NiO+10%YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>5</td>
<td>10%Pr$<em>{0.3}$Ce$</em>{0.35}$Zr$_{0.35}$O$_2$+90%NiO/YSZ</td>
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</tr>
<tr>
<td>6</td>
<td>1%Ru/10%Pr$<em>{0.3}$Ce$</em>{0.35}$Zr$_{0.35}$O$_2$+90%NiO/YSZ</td>
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</tr>
<tr>
<td>7</td>
<td>10%Pr$<em>{0.25}$Ce$</em>{0.5}$Zr$_{0.25}$O$_2$+90%NiO/YSZ</td>
<td>0</td>
</tr>
<tr>
<td>8</td>
<td>1%Ru/10%Pr$<em>{0.25}$Ce$</em>{0.5}$Zr$_{0.25}$O$_2$+90%NiO/YSZ</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>10%Pr$<em>{0.15}$La$</em>{0.15}$Ce$<em>{0.35}$Zr$</em>{0.35}$O$_2$+90%NiO/YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>10</td>
<td>1%Ru/10%Pr$<em>{0.15}$La$</em>{0.15}$Ce$<em>{0.35}$Zr$</em>{0.35}$O$_2$+90%NiO/YSZ</td>
<td>0.3</td>
</tr>
<tr>
<td>11</td>
<td>10%Pr$<em>{0.15}$La$</em>{0.15}$Ce$<em>{0.5}$Zr$</em>{0.2}$O$_2$+90%NiO/YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>12</td>
<td>Ru/10%Pr$<em>{0.15}$Ce$</em>{0.5}$Zr$_{0.2}$O$_2$+90%NiO/YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>13</td>
<td>10%Pr$<em>{0.15}$Sm$</em>{0.15}$Ce$<em>{0.35}$Zr$</em>{0.35}$O$_2$+90%NiO/YSZ</td>
<td>0.1</td>
</tr>
<tr>
<td>14</td>
<td>Ru/10%Pr$<em>{0.15}$Sm$</em>{0.15}$Ce$<em>{0.35}$Zr$</em>{0.35}$O$_2$+90%NiO/YSZ</td>
<td>0</td>
</tr>
<tr>
<td>15</td>
<td>10%Pr$<em>{0.15}$Sm$</em>{0.15}$Ce$<em>{0.5}$Zr$</em>{0.2}$O$_2$+90%NiO/YSZ</td>
<td>0</td>
</tr>
<tr>
<td>16</td>
<td>Ru/10%Pr$<em>{0.15}$Sm$</em>{0.15}$Ce$<em>{0.5}$Zr$</em>{0.2}$O$_2$+90%NiO/YSZ</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 2. Concentrations of products (H$_2$, CO and CO$_2$) in the ethanol steam reforming at 700 °C for different composites prepared by modified Pechini method (1-4) and by impregnation (combinatorial) rout (4-16). Contact time 0.036s, feed composition 0.5% C$_2$H$_5$OH + 2.5% H$_2$O in He.
Fig. 20. Rates of products formation in ethanol steam reforming at 700 °C on nanocomposites. Contact time 36ms, feed 0.5% EtOH + 2.4%H\textsubscript{2}O in He. 1-10%SmPrCeZr+55%NiO+35%YSZ, 2-80% SmPrCeZr +10%NiO +10%YSZ, 3-50%LaMnCrPr+30%NiO+20%YSZ, 4-80%LaMnCrPr +10%NiO+10%YSZ

Some variation of performance with nanocomposites composition and preparation route is worth commenting. For samples based on composite III prepared via successive impregnation, the lowest performance was demonstrated by nanocomposites containing Pr\textsubscript{0.15}La\textsubscript{0.15}Ce\textsubscript{0.7-x}Zr\textsubscript{x}O\textsubscript{2} (Table 2). This clearly correlates with the highest temperatures of H\textsubscript{2} TPR peaks for these samples (Sadykov et al., 2009b) reflecting their lower reducibility, and, hence, a lower mobility and reactivity of lattice oxygen due to stabilizing effect of a big basic La cation. Since in diluted ethanol+ water feeds the catalysts were tested after pretreatment in O\textsubscript{2}, this correlation can be explained either by insufficient reduction of Ni in La-containing catalysts, or by a lower mobility and reactivity of surface hydroxyls/hydroxocarbonates bound with big La cation. The latter can result in the surface coking decreasing performance. Indeed, for nanocomposite promoted by Pr\textsubscript{0.15}La\textsubscript{0.15}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2} with the lowest hydrogen concentration in products (Table 2), TPO oxidation after reaction revealed accumulation of up to 18 monolayers of carbonaceous deposits on the surface.

Effect of supporting Ru on promoted composite III strongly depends upon the composition of complex oxides (Table 2). For PrSmCeZr- promoted composites possessing a high activity, supporting Ru slightly decreases hydrogen yield. For composite promoted by Pr\textsubscript{0.15}Sm\textsubscript{0.15}Ce\textsubscript{0.35}Zr\textsubscript{0.35}O\textsubscript{2}, co-promotion with Ru increases CO\textsubscript{2} yield but decreased CO and H\textsubscript{2} yield. Hence, at short contact times and in diluted feeds, water gas shift reaction is apparently not equilibrated.

When Ru was supported on less active composites promoted by La-containing oxides, another effects were observed: both hydrogen and CO\textsubscript{x} yields were increased. While the increase of CO yield can be assigned to acceleration of reverse water gas shift reaction catalyzed by Ru particles weakly interacting with promoting fluorite-like oxides in these composites (vide supra), increase of hydrogen and CO\textsubscript{2} content in the feed suggests that the overall transformation of ethanol along the steam reforming route is accelerated as well, perhaps, due to decreasing the surface coking and/or accelerating other stages of intermediates (acetate etc) transformation into hydrogen and carbon oxides.

For SmPrCeZrO-promoted nanocomposites prepared via one-pot Pechini route (sample 1, Table 2) performance is lower than for nanocomposites of the same composition prepared via consecutive impregnation (sample 13, Table 2). This demonstrates that nanocomposite
microstructure affected by the preparation route plays important role in performance of these catalysts in steam reforming of ethanol. The decrease of Ni content in nanocomposite from 55 to 10% only moderately decreased performance (Table 2), which can be explained by a higher dispersion of Ni (vide supra). Nanocomposites promoted by LaPrMnCrO perovskite and prepared via Pechini route demonstrate a higher activity in ethanol steam reforming than catalysts promoted by doped ceria-zirconia oxides and prepared by the same method (Table 2, Fig. 20). Since the trend was usually reversed for CH₄ steam reforming, such a simple explanation as a higher dispersion of Ni more easily incorporated into the perovskite structure could be not sufficient. Perhaps, for ethanol steam reforming presence of red-ox cations such as Mn which can be involved in H₂O activation along with Ni (vide supra) can be important as well.  

4.3 Catalytic performance of nanocomposites supported on structured substrates  

For supporting thin layers of nanocomposites, Ni/YSZ anode plates, Ni-Al foam substrates as well as thin-foil or gauze FechrAlloy substrates were used. The FechrAlloy gauze substrate (woven from the wires with diameter 0.2 mm and ca. 0.2 mm spacing) was first precovered by a thin (ca. 5–10 microns) corundum sublayer by blast dusting technique (Ulyanitskii et al., 2006) followed by washcoating with La-stabilized γ-Al₂O₃ (3.6 wt.%) from an appropriate suspension followed by calcination under air at 1100ºC for 2 h. Before supporting active components, the gauze was cut into square pieces to be stacked with Ni-Al foam plates. Composite powders synthesized via one-pot Pechini route (vide supra) were ultrasonically dispersed in isopropyl alcohol with addition of polyvinyl butyral as a binder to make a slurry. Thin layers of composites were supported on heat-conducting substrates using these slurries and slip casting or painting procedures followed by drying and calcination at 1100ºC after each supporting step until loading of 4-7 wt.% was achieved. Ru or Pt were supported by the incipient wetness impregnation followed by drying and calcination under air at 800ºC. Catalytic performance of monolithic catalysts with nanocomposite active components on different substrates in the reaction of natural gas and ethanol steam reforming in concentrated feeds was studied in the stainless steel flow reactors equipped with external heating coils (Sadykov et al., 2010a) using a pilot-scale installation (feed rate up to 1 m³/h). Water and ethanol were supplied by a pump and sprayed via a nozzle into a specially designed monolithic FechrAlloy honeycomb evaporation/mixing unit heated by passing the electric current (Sadykov et al., 2009a). Reagents and products concentrations were analyzed by GC. Fig. 21 compares methane conversions for different plate-like structural elements in stoichiometric steam/methane feed with supported layers of nanocomposite containing fluorite-like oxide promoter and Ru. While at temperatures below 550ºC the initial Ni/YSZ anode substrate is not active, supporting composite layers on all substrates provides a reasonable activity in the low-temperature range. Activity of these layers on different substrates is comparable, which agrees with reasonably high intrinsic activity of nanocomposite active components (vide supra). This allows to use these structural elements for in-cell steam reforming of methane to provide an efficient heat management. A high activity of unpromoted Ni/YSZ anode platelet manufactured by the Research Center of Jülich (Germany) at temperatures exceeding 600ºC is worth noting as well. In this case, as judged by the carbon balance, a slow coke accumulation in stoichiometric feed takes place which is to deactivate anode sooner or later.
When nanocomposite layers supported on different substrates contain as co-promoters Ru and perovskite-like oxide, performance is better than for combination of Ru and fluorite-like oxide (Fig. 21), following similar trend for powdered samples (vide supra). Combination of Pt with fluorite-like oxide as co-promoters in supported nanocomposite layers on different substrates also provides a high and stable activity in methane steam reforming even in feeds with higher (20%) concentration of methane close to realistic composition (Fig. 22).

Fig. 21. Temperature dependence of CH$_4$ conversion on catalytic platelets of 1x2 cm$^2$ size. 8% CH$_4$ +8% H$_2$O in He, contact time 25 ms. Left: 1%Ru/Pr$_{0.3}$Ce$_{0.35}$Zr$_{0.35}$O$_2$/Ni/YSZ layers on Ni/YSZ anode platelet (1) foam Ni-Al substrate (2) and Crofer interconnect (3). 4-unpromoted Ni/YSZ anode platelet. Right: 0.5%Ru/La$_{0.8}$Pr$_{0.2}$Mn$_{0.2}$Cr$_{0.8}$O$_3$/Ni/YSZ (1) and 0.5%Ru/Pr$_{0.3}$Ce$_{0.35}$Zr$_{0.35}$O$_2$/Ni/YSZ (2) layers on Ni/YSZ anode substrate.

As follows from both optical and SEM images after stability tests (Fig. 23), supported nanocomposite layer retains its integrity without any cracks or detachment from the underlying anode substrate despite transformation of NiO into Ni in composite as well as in the anode substrate in the reaction conditions. This is provided by a close composition of both substrate and supported layers ensuring matching of chemical and thermal coefficients of shrinkage/expansion. Developed porosity of supported layers helps to avoid diffusion limitations for rather fast CH$_4$ SR reaction.

Fig. 22. Conversion of CH$_4$ (1,2) and H$_2$ concentration in converted dry feed (3,4) for CH$_4$ steam reforming on nanocomposite Pt/Pr$_{0.3}$Ce$_{0.35}$Zr$_{0.35}$O$_2$/Ni/YSZ layers supported on one side of Ni/YSZ anode platelet (1,3) or porous Ni-Al substrate (2,4). CH$_4$/H$_2$/Ar = 20/20/60. Contact time 25 ms.
In addition to the direct in-cell steam reforming of methane occurring on catalytically active anodes of SOFC, nanocomposite catalysts supported on heat-conducting substrates such as Crofer interconnects or foam Ni-Al alloy can be efficiently applied for so called indirect in-cell steam reforming of methane within the anode compartment of a stack (in fuel channels etc) (Dicks, 1998). In this case, the effect of the oxygen transfer across the cell under load is of much less importance, so selection of active components can be primary oriented on achieving the highest performance in feeds with steam/methane ratio close to stoichiometry and its stability.

For the indirect in-cell reforming of methane on separate structural elements comprised of nanocomposite active components supported on heat –conducting metal alloy substrates, the problem of thermal expansion compatibility with substrates is less crucial as compared with that for porous nanocomposite layers on anode platelets. Hence, content of Ni in active component can be decreased to improve its resistance to sintering and coking. On the other side, decreasing Ni content could decrease catalytic activity, especially in feeds with the water excess (vide supra).

Systematic studies of the effect of Ni content in nanocomposite active component on their performance revealed that the best compromise is achieved at Ni content in the nanocomposite with YSZ and complex promoting perovskite-like or fluorite-like oxides around 25% (NiO content in green nanocomposite 30 wt.%). As demonstrated in Fig. 24, in this case the active component provides required level of activity and on-stream stability when tested both as a fraction and as a porous layer supported onto foam Ni-Al substrate. Note that the temperature dependence of conversion is practically identical for the fraction and platelet. Hence, mass and heat transfer effects are indeed negligible for this type of substrate.

To check the effect of up-scaling the size of structured catalysts on their performance in the reaction of methane steam reforming, a package comprised of stacked foam platelets and gauzes was tested in concentrated natural gas/steam feeds.

For the stack comprised of either of 3 Ni-Al foam plates (Fig. 25) or 12 Ni-Al-foam plates and 11 sheets of Fecralloy gauzes loaded with La_{0.8}Pr_{0.2}Mn_{0.2}Cr_{0.8}O_3 + NiO + YSZ + Ru active component (Fig. 26), hydrogen concentration in the effluent was nearly identical at the same temperature and contact time. This means that heat and mass transfer does not affect strongly performance of these structured catalysts, which is rather good providing >45% \( \text{H}_2 \) in the effluent.
Fig. 24. Temperature dependence of CH$_4$ conversion (left) and its variation with time-on-stream at 600 ºC (right) for 0.25 mm fraction of nanocomposite V +0.7Ru (1), and 1x2 cm$^2$ Ni-Al platelet with supported active component (2, 3). Loading 5.3 wt. % nanocomposite V +0.85 wt.% Ru (2) or 3.8 wt. % nanocomposite V +0.47 wt.% Ru (3). Contact time 10 ms for (1), 25 ms for (2, 3) and 50 ms for testing stability with time-on-stream. Feed composition 20% CH$_4$ +40% H$_2$O, Ar balance.

Fig. 25. Effect of temperature (left) and contact time (STP) at 600ºC (right) on concentration of components in effluent for steam reforming of natural gas on stack of 3 plates 40x40x1 mm$^3$ each. Feed composition 35% of natural gas + 60% H$_2$O + 5% N$_2$; contact time 0.15 s at 600ºC or 0.5 s at STP (left).

Tests for 100 h with start-up and shut-down of pilot installation each day (8 hours working time per day) confirmed stability of this level of H$_2$ content in effluent (Fig. 26). The same package of 12 Ni-Al-foam plates and 11 sheets of Fecralloy gauzes loaded with La$_{0.8}$Pr$_{0.2}$Mn$_{0.2}$Cr$_{0.8}$O$_3$ + NiO + YSZ + Ru active component was used for oxysteam reforming of ethanol in pilot installation. In this process performance was also stable providing a high concentration of hydrogen (Fig. 27). The increase of steam excess in the feed helps to increase the hydrogen yield by increasing ethanol conversion as well as by decreasing content of such by-products as methane and ethylene.
5. Kinetic analysis of methane steam reforming on anode structural elements

The kinetics of the methane–steam reaction is based upon a 13-step mechanism of $\text{H}_2$; CO and $\text{CO}_2$ formation with three rate determining steps (Xu & Froment, 1989).

Three reversible macrosteps are as follows:

- **Steam reforming (SR)**

  \[
  \text{CH}_4 + \text{H}_2\text{O} \leftrightarrow 3\text{H}_2 + \text{CO} \quad \Delta_h^{298K} = 206.2 \text{ kJ mol}^{-1} \tag{1}
  \]
• Water-gas shift (WGS)

\[
\text{CO} + \text{H}_2\text{O} \leftrightarrow \text{CO}_2 + \text{H}_2 \quad \Delta_r H_{298 K} = -41.2 \text{ kJ mol}^{-1}
\] (2)

• Complex shift reaction (SSR)

\[
\text{CH}_4 + 2\text{H}_2\text{O} \leftrightarrow 4\text{H}_2 + \text{CO}_2 \quad \Delta_r H_{298 K} = 165 \text{ kJ mol}^{-1}
\] (3)

Kinetic parameters of these steps were estimated by analyses of experimental data obtained in a slot-like reactor for the nanocomposite platelet with size 1x2 cm² loaded with 5.3% wt.% of nanocomposite 50%LaMnPr+30%NiO+20% YSZ promoted by 0.85 wt.% Ru. The power law rate equations for the reactions (1) - (3) were expressed in the general form:

\[
R_{r,j} \left[ \text{kg mol} \cdot \text{m}^{-3} \cdot \text{s} \right] = k_{f,r} \prod_{j=1}^{N_r} \left[ C_{j,r} \right]^{\eta_{j,r}} - k_{b,r} \prod_{j=1}^{N_r} \left[ C_{j,r} \right]^{\eta_{j,r}},
\]

where Arhenius parameters for forward reaction rate constants are written in the form:

\[
k_{f,r} = A_r T^{E_r} \exp \left( -\frac{E_r}{RT} \right).
\]

Here, \(N_r\) is number of chemical species in the overall molecular reaction \(r\) (SR, WGS, and SSR); \(C_{j,r}\) is molar concentration of each reactant and product species \(j\) in reaction \(r\) (kg mol/m³); \(\eta_{j,r}\) is forward rate exponent for each reactant and product species \(j\) in reaction \(r\); \(\eta_{j,r}\) is backward rate exponent for each reactant and product species \(j\) in reaction \(r\). The backward rate constant for reaction \(r\), \(k_{b,r}\), were computed from the forward rate constant using the following relation: \(k_{b,r} = \frac{k_{f,r}}{K_{eq,r}}\), where \(K_{eq,r}\) is the equilibrium constants for the three main reactions (1)- (3) respectively:

\[
K_{eq,SR} \left[ \text{atm}^2 \right] = \frac{P_{\text{CO}P_{\text{H}_2}^3}}{P_{\text{CH}_4}P_{\text{H}_2\text{O}}^3} = \exp \left( -\frac{\Delta G_{SR}}{RT} \right),
\]

\[
K_{eq,WGS} \left[ \text{atm}^0 \right] = \frac{P_{\text{CO}_2P_{\text{H}_2}^2}}{P_{\text{CH}_4}P_{\text{H}_2\text{O}}} = \exp \left( -\frac{\Delta G_{WGS}}{RT} \right),
\]

\[
K_{eq,SSR} \left[ \text{atm}^2 \right] = \frac{P_{\text{CO}_2P_{\text{H}_2}^2}}{P_{\text{CH}_4}P_{\text{H}_2\text{O}}^3} = \exp \left( -\frac{\Delta G_{SSR}}{RT} \right).
\]

Respective values of effective kinetic parameters for the reactions (1) - (3) obtained by fitting the experimental data are given in Table 3. The fixed conversion temperature dependences were analyzed with the developed rate equations (Table 3).
Table 3. Kinetic parameters of the rates of three basic macrosteps for the catalytic platelet loaded with nanocomposite active component.

To perform numerical experiments a steady state plug-flow reactor model describing the change of each reaction component along the catalytic platelet is written in the differential form:

\[
\frac{d (uc_i)}{dl} = S_{sp} \sum_{j=1}^{3} v_{ij} r_j
\]  

Here, \( c_i \) is concentration of i-component in mol/m\(^3\), \( u \) - superficial velocity of gas mixture through the reactor (m/s), \( S_{sp} \) - specific surface area, (m\(^{-1}\)), \( v_{ij} \) - stoichiometric coefficients; \( r_j \) - reaction rate for the three global reactions (mol/m\(^2\)*s\(^{-1}\)), expressed in the empirical form of the power law rate kinetics for the global reactions (1)-(3) as as 

\[
r_j = \frac{R_j}{S^*}, \quad \text{where} \quad S^* \quad \text{active surface area (m}^2/\text{kg}).
\]

A second-order Rozenbroke algorithm with an automatic choice of the integration interval was used to solve the set of equations. The Fortran computed code was developed to implement the described algorithm. The predictions based on the kinetic models with the parameters summarised in Table 3 were compared with the experimental data (Fig. 28). A good agreement demonstrates that the power law model is able to predict with certainty the reaction behavior within the temperature region 600-700°C, which is of special interest in the case of in-cell methane reforming for intermediate temperature solid oxide fuel cells.

The same model was applied to analysis of experimental data on the natural gas steam reforming over stack comprised of three parallel Ni-Al platelets (50x50x1 mm) loaded with the same nanocomposite active component and separated by 1 mm gaps. In these experiments operational temperature, the feed linear velocity and the steam-to-natural gas ratio were varied. A comparison of simulated and experimental data obtained for this stack has been performed by using kinetic parameters estimated for one small platelet (Table 3). The general quality of results obtained is illustrated in Figures 29 (a,b). A good agreement between experimental and simulated data demonstrates that the simple plug-flow reactor model is able to describe the reaction behaviour in the wide range of the experimental conditions without taking into account heat and mass transfer effects due to high thermal conductivity and developed macroporosity of Ni-Al foam substrates.
Fig. 28. Experimental and simulated (with kinetics – solid lines, thermodynamic predictions – dash lines) data (dry basis) on the effect of the temperature on the methane steam reforming reaction over Ni-Al foam platelet loaded with 5.3% wt.% of 50% LaMnPr+30%NiO+20% YSZ nanocomposite promoted by 0.85% Ru.

Fig. 29. Effect of the operational temperature (a) and linear velocity of the feed gas (STP) (b) on the concentration of components in the product gas. Symbols – experimental data, lines – numerical experiments.

6. Results of catalytic element testing in planar SOFC

The cell tests were performed in the Energy Research Center (Petten, Netherlands) in a test bench with an alumina housing for 5x5 cm² cells (Ouweltjes et al., 2008). Fuel cells were assembled in configuration Pt grid/LSC/20GCO/8YSZ/Ni-YSZ anode substrate/Ni-Al foam catalytic plate/Ni grid. Here platinum grid was used for cathode current collection, LSC is cathode La-Sr-cobaltite layer, 20GDC- Ce₀.₈Gd₀.₂O₂₋y interlayer between cathode and a thin layer of YSZ supported on a planar Ni-YSZ anode substrate. Nickel mesh as anode current collector was pressed to Ni-Al catalytic plate, so current was passed through it. A stream of CH₄ + H₂O feed (H₂O/CH₄ =2) or humidified H₂ was passed along the catalytic plate loaded with 5 wt.% of nanocomposite 50% LaMnCrPr + 30% NiO +
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20% YSZ promoted by 1.3 wt.% Ru. Air stream was used as oxidant from the cathode side. Electrochemical characterisation was performed by V/I curves and impedance spectroscopy with a Solartron 1255/1287 set-up. At 600 °C and 700 mV the area specific resistance (ASR) was estimated to be 0.77 Ohm cm² without any contribution of catalyst to $R_{\text{ohm}}$, which meets target of operation for intermediate temperature solid oxide fuel cells.

Fig. 30 presents I- V characteristics of this cell with CH₄ or H₂ as a fuel. As follows from these data, at 800 °C cell performance for both types of fuels is comparable, which demonstrates a high efficiency of planar catalytic element in steam reforming of methane. Power density up to 350-850 mW/cm² is achieved in 600-800 °C range, which is promising for the practical application. At CH₄ flow 90 ml/min and 600 °C CH₄ conversion increases from 47 to 55% with increasing current from 0 to 4A. This conversion is rather close to values obtained at this temperature for single platelet with this active component (Fig. 21).

Fig. 30. I- V characteristics of cell equipped with Ni-Al foam-supported nanocomposite catalyst using CH₄ or H₂ as a fuel.

7. Conclusion

Nanocomposite materials developed in this work based upon Ni and doped zirconia electrolyte allow to provide efficient and stable in-cell steam reforming of methane and ethanol required for IT SOFC as well as design of monolithic catalysts of transformation of biofuels into syngas. This is provided by optimization of their composition and preparation procedures ensuring developed interfaces between components activating fuel molecules (Ni, Ni-Pt/Ru alloys) and water molecules (complex oxides with perovskite and fluorite structures). Performance of best compositions supported as porous strongly adhering layers on anode cermets platelets, FeCrAlloy gauzes and porous Ni-Al foam substrates was estimated as well and demonstrated to be high and stable to meet target of internal reforming of fuels in the intermediate temperature solid oxide fuel cells. No cracking or detachment of layers after reaction was observed. Analysis of methane reforming kinetics catalyzed by structured catalytic elements with supported nanocomposite layers has been carried out. Performance of catalytic plates in IR mode of ECN cell meets target of design of
solid oxide fuel cells with internal reforming of methane by area specific resistance, activity and power density.

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Advances in Nanocomposites - Synthesis, Characterization and Industrial Applications was conceived as a comprehensive reference volume on various aspects of functional nanocomposites for engineering technologies. The term functional nanocomposites signifies a wide area of polymer/material science and engineering, involving the design, synthesis and study of nanocomposites of increasing structural sophistication and complexity useful for a wide range of chemical, physicochemical and biological/biomedical processes. "Emerging technologies" are also broadly understood to include new technological developments, beginning at the forefront of conventional industrial practices and extending into anticipated and speculative industries of the future. The scope of the present book on nanocomposites and applications extends far beyond emerging technologies. This book presents 40 chapters organized in four parts systematically providing a wealth of new ideas in design, synthesis and study of sophisticated nanocomposite structures.

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